## PATENT SPECIFICATION

NO DRAWINGS

1,117,129



Inventors:-SABBAT JOHN STRIANSE and MARK ROY HAVASS.

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Int. CL:-A 61 k 7/06.

## COMPLETE SPECIFICATION

## A Cosmetic Preparation

We, YARDLEY AND COMPANY LIMITED, a
British Company of London, England, do
hereby declare the invention, for which we
pray that a patent may be granted to us,
and the method by which it is to be performed, to be particularly described in and
by the following statement:

This invention relates to hair grooming compositions, including hair fixature com-

10 positions.

For many years compositions based on hydrocarbon oils and oil gels, such as petrolatum, have been sold for restoring oil to dry hair and scalp, to brighten the hair and to impart to it a measure of manageability. Such compositions suffer from a variety of

disadvantages.

The oil gel types of hair grooming compositions, often called "pomades" or "solid brilliantines" are most used for hair control; but they have poor lubricity, and poor spreadability, and thus are likely to leave a high degree of stickiness or greasiness on the hair, which is objectionable to many persons. Such compositions made by bodying mineral oil by use of known agents such as aluminium stearate or paraffin wax, are subject to separation of liquid oil (syneresis), friability of the gel structure, shrinkage within the jar, and poor spreadability. These characteristics have made such compositions less than satisfactory and have limited their

The heavier oils in liquid form also are difficult to spread thinly and uniformly on the hair; whereas the thinner oils tend to run off the hands and hair and down onto one's face or neck or along one's wrist, onto sleeves or other parts of one's clothing.

The liquid oil compositions are also less than satisfactory in their grooming and fixative effects. The oil is not absorbed by the hair and, to the extent that it provides hair

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control, it also causes an objectionable oiliness and apparent matting, "plastering 45 down" of the hair.

The present invention is based upon the discovery that desirable and effective hair grooming compositions can be made by at least partly dissolving polyamide material in 50 a liquid, oily, non-polar solvent consisting of or including a substance having a chain of at least 10 carbon atoms in its molecule, the polyamide material being a reaction product of an aliphatic poly-carboxylic acid and an 55 alkylene polyamine and having an average molecular weight between 2000 and 14000. Advantageously the polyamide material is of the type set forth in U.S. Patents Nos. 2,450,940 and 2,379,413, having an average molecular weight between 2000 and 10000 and being reaction products of aliphatic dicarboxylic acids and di- or polyamino compounds.

The present invention provides a hair 65 grooming and hair treating composition which is free from the serious disadvantages of the prior compositions, and thus provides improvements in such compositions and in the art of hair grooming which have long 70 been sought, but had seemed unartainable.

The composition may include a cosolvent as well as the oily vehicle (the solvent). The cosolvent dissolves the resin and is miscible with the oil so as to bring the composition 75 into the form of a stable gel or suitably bodied or polymerized liquid. Other ingredients may be included for example, ordinary cosmetic diluents and ingredients, for example, scents and tinting colours.

These may be added to the oily vehicle non-polar materials solid at ambient temperature. The oily vehicle may include, for example, mineral oils which are advantageously of 50-70 Saybolt viscosity. Higher 55 viscosity oils have less compatibility with

[Price 4s. 6d.]

Price 33p

Price Top

the polyamide material and give less clear

The cosolvents or coupling agents chosen cosmetically acceptable compounds 5 which form with the resin a solution which ATE is miscible with the oily vehicle. In general, the cosolvent is a substance in which both the resin and the oil are soluble. The most advantageous liquids as cosolvents are of the 10 class of fatty acids, alcohols and glycol esters having a hydrocarbon radical of the kind found in vegetable oils, most advantageously the hydrocarbon radical has a straight chain of 12-18 carbon atoms.

15 Typical examples of cosolvents are oleic acid, linoleic acid, mixtures of oleic and linoleic acids, diethanolamine linoleate. oleyi alcohol, propylene giycol mono laurate, propylene glycol di-laurate, propylene glycol monomyristate, propylene glycol mono-oleate, lauryl lactate, myristyl lactate, methyl salicylate, castor oil, ethanol, isopropanol. di-butyl phthalate, di-butyl sebacate, and dioctyl sebacate, or mixtures thereof. How-25 ever, some of the above cosolvents, for example propylene glycol mono laurate, can be used without the oily vehicle in which case they act as the oily liquid non-polar solvent.

30) The polyamide material as already indicated above, is advantageously a solid resinous, condensation product of an aliphatic dicarboxylic acid and a diamine (inclusive of compounds having at least one 35 alkylene and at least two amino groups, respectively) soluble in at least some organic

Suitable resins of this type are solvents. available commercially from General Mills, Inc., under the name "Versamid" (Registered Trade Mark), and from Olin-Mathison 40. Chemical Corp., under the name "Omamid" (Registered Trade Mark), for example Omamid "S" or Omamid "C". They are tough thermoplastic resins of the polyamide type insoluble in water and in many ketones 45 and ester solvents such as carboxylic acid amides, alcohols and chlorinated hydrocarbons, depending upon the particular acids and amines which have been used to form the resin and also upon the polymer length. 50 Monohydric alcohols, especially those having 3-8 carbon atoms, and chlorinated hydrocarbons are generally effective and hy-drocarbons and ether solvents are in some cases effective per se and in some cases 55 effective only in mixtures with other solvents. Specifically, such solvents include namyl alcohol, iso amyl alcohol, benzene, iso butyl alcohol, ethyl alcohol, n-octyl alcohol, mono butyl ether of ethylene glycol, mono 60 ethyl ether of ethylene glycol, n-propyl al-cohol, iso propyl alcohol, turpentine, xylene and mixtures thereof. Chloroform, methylene chloride, turpentine and xylene, even though effective for solvent purposes, are not 65 recommended for hair grooming cosmetica because of odour. In general, solubility is low or absent with simple hydrocarbons, but as indicated above, they may be useful in mixtures with other solvents.

Other properties of these resins appear as

Omamid

follows: --

				v ersumua			•	
75	Resin type Specific gravity †Colour, Gardner Melting point °C	900 .98 12 180-190	930 .98 12 105-115	940 .98 12 105-115	950 .98 12 43-55	100 .98 12 43-55	<i>C</i> .9799 12- 14	.9496 11- 12
	Softening point °C (ring and ball) Viscosity						90-100	100-120
80	†Brookfield at 150°C Solid No. 2 Spindle		30-45	15-30	7-15	10-15	32- 52 12	55- 75 12
	Acid value Amine value*	3	3	3	3	83-93		 emine

\*Amine value is the weight of KOH, in milligrams, equivalent to the free amine groups in one gram of the resin.

†Gardner Colour Scale (Transparent Liquids) Ref: American Society for Testing of Materials (Standards) Part 21, 1964.

†Brookfield viscometer data, Ref: American Society for Testing of Materials (Standards), Part 26, 1964.

90 "Versamid" polyamide resins are thermoplastic condensation products of polymerized linoleic acid with various polyamine compounds such as ethylene diamine, and diethylene triamine. Resins of average 95 molecular weights of 5000-10000 have been found best for the present invention. These resins are commercially available in hard, brittle resin (No. 900) of melting point 180-190°C, tough flexible resins (No. 930 and 100 940) melting points 105-115°C, and in semi-

solid, soft tacky resin (No. 100), melting point 43-55°C., and with some wax added (No. 950), which results in some turbidity in the final product. These resins are compatible with each other so that by blending 105 them almost any desired properties in the aforesaid melting point range can be attained.

The tendency to syneresis of the oilpolyamide-cosolvent gels can be controlled 110

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by use of amides having 12 to 18 carbon	2. Simple system for clear gel:		
atoms in their molecules, for example di-	Polyamide 8000 average MW	5.00	
ethanolamides fulfilling this condition and/	Propylene glycol mono laurate Light mineral oil	70.00 25.00	
or by curing the gels by holding them at 5 temperatures between their melting point	Light immeration	25.00	70
and the ambient temperature.		100.00	70
Hair treating gels are prepared by dis-	3. Simple system for cloudy gel:	100.00	
solving the polyamide resins in the hot or-	Polyamide 8000 MW (average)	5.00	
ganic system comprising the oily vehicle and	Propylene glycol mono laurate	47.50	
10 the cosolvents (if included). Upon cooling,	Light mineral oil	47.50	75
a gel structure is produced, and the pro-			. •
perties will vary depending upon the amount		100.00	
of resin employed, the composition and	4. Simple system including large		
molecular weight of the resin, and the com-	percentage of mineral oil		
15 patibility and solubility of the resin in the	(cloudy soft gel):		80
vehicle chosen. The solubility of the poly-	Polyamide 8000 MW (average)	2.00	
amide resin in preferred solvent systems in-	Oleic Acid	13.00	
creases with temperature. Whenever the	Light mineral oil	85.00	
solubility limits of the polyamide resin in a		100.00	
21) particular solvent system are exceeded, a gel	Cimple system for clear sel	100.00	85
results which is thermally and mechanically	5. Simple system for clear gel using a blend of polyamide		
reversible. Gels can be produced of a con- sistency from a soft jelly-like to a firm rigid	resins:		
structure, or of grainy, "crystal-like" struc-	Polyamide 8000 MW (average)	2.50	
25 ture or an amorphous, smooth glass-like	Polyamide 5000 MW (average)	2.50	<b>90</b> .
structure.	Propylene glycol mono laurate	70.00	
Mixing at high temperatures, e.g. in the	Light mineral oil	25.00	
range 100-115°C may cause crystallization,	•		
whereas if the composition is mixed below		100.00	
30 100°C and poured into moulds or jars at	The light mineral oil referred		
about 55°C and then kept for a substantial	example is Marcol GX available		
curing time, e.g. 1-6 days at an intermediate	Standard Oil Co. The use of oth		
temperature, e.g. 37°C±2°C., a smoother	oils in many systems produces		
structure and freedom from syneresis are	cloudy gels. However, these other oils can be made to give crystal		
35 attained. (See Example 9 below). Such compositions are economical, thixo-	tems by rebalancing the cosolvent		- 100
tropic and less subject to syneresis than the	The gels of Examples 2 to 5 ma		a
brilliantine type compositions having a metal	by heating the oily solvent and the		
soap, such as aluminium stearate, as the	to slightly above the melting po		
40 gelling agent, in mineral oil.	polyamide which is then introduce	ed into the	e 105
It is an important advantage of the inven-	mixed solvents with agitation unti	l the mas	3
tion that crystal clear gels can be made, al-	is homogeneous. The temperate	ure is re	-
though opaque or cloudy gels are also with-	duced to below 100°C and variou		
in the broader scope of the invention. Such	are then introduced into the mass		
45 gels, whether crystal clear or non-clear, pro-	ring. It is then cured and filled in	to smorpi	c 110
vide a thixotropic, non-greasy solid gel,	A further improvement has been	en attaine	đ
which is liquefied quickly to a fluid by rub- bing and is thus readily spread on and	by using appropriate mixtures of		
throughout the hair leaving the hair con-	of the resin and mineral oil. This		
50 trolled and well groomed but with a natural	crease of the mineral oil content		
soft appearance free from the "plastered-	sponding decrease in cost, while		
down" look and greasiness which have made	higher temperature stability and	clarity o	ď
most hair dressing compositions unaccept-	the gel:		
able to many persons.	Example 6 — A clear gel system	using tw	0
55 Following are examples of compositions	cosolvents:	D	120
embodying the present invention which are		Percentag	
suitable for hair grooming products:—	Dolumnida 2000 amana MI	by weigh	
Examples of Gels % By Weight	Polyamide 8000 average MW Propylene glycol mono laurate	5.00 19.10	
1. Clear tacky and substantially	Oleic Acid	10.90	125
60 free of syneresis Polyamide 5000-8000 average	Light mineral oil	64.00	
MW 5.00	Perfume	1.00	
Castor Oil 95.00			
	•	100.00	
65 100.00			130
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by use of amides having 12 to 18 carbon	2. Simple system for clear gel:		
atoms in their molecules, for example di-	Polyamide 8000 average MW	5.00	
ethanolamides fulfilling this condition and/	Propylene glycol mono laurate	70.00	
or by curing the gels by holding them at	Light mineral oil	25.00	
5 temperatures between their melting point			70
and the ambient temperature.		100.00	
Hair treating gels are prepared by dis-	3. Simple system for cloudy gel:		
solving the polyamide resins in the hot or-	Polyamide 8000 MW (average)	5.00	
ganic system comprising the oily vehicle and	Propylene glycol mono laurate	47.50	
10 the cosolvents (if included). Upon cooling,	Light mineral oil	47.50	75
a gel structure is produced, and the pro-			
perties will vary depending upon the amount		100.00	
of resin employed, the composition and	4. Simple system including large		
molecular weight of the resin, and the com-	percentage of mineral oil		
15 patibility and solubility of the resin in the	(cloudy soft gal):		80
vehicle chosen. The solubility of the poly-	Polyamide 8000 MW (average)	2.00	
amide resin in preferred solvent systems in-	Oleic Acid	13.00	
creases with temperature. Whenever the	Light mineral oil	85.00	
solubility limits of the polyamide resin in a			
21) particular solvent system are exceeded, a gel		100.00	85
results which is thermally and mechanically	5. Simple system for clear gel		
reversible. Gels can be produced of a con-	using a blend of polyamide		
sistency from a soft jelly-like to a firm rigid	resins:		
structure, or of grainy, "crystal-like" struc-	Polyamide 8000 MW (average)	2.50	
25 ture or an amorphous, smooth glass-like	Polyamide 5000 MW (average)	2.50	<b>90</b> .
structure.	Propylene glycol mono laurate	70.00	
Mixing at high temperatures, e.g. in the	Light mineral oil	25.00	
range 100-115°C may cause crystallization,			
whereas if the composition is mixed below		100.00	
30 100°C and poured into moulds or jars at	The light mineral oil referred	to in thi	s 9\$
about 55°C and then kept for a substantial	example is Marcol GX available	from Eas	0
curing time, e.g. 1-6 days at an intermediate	Standard Oil Co. The use of other		
temperature, e.g. 37°C±2°C., a smoother	oils in many systems produce	s hazy t	0
structure and freedom from syneresis are	cloudy gais. However, these off		
35 attained. (See Example 9 below).	alle one he made to sime service	Alees are	_ 1110
	oils can be made to give crystal	. cacan sy	- 144
	tems by rebalancing the cosolven	t fraction.	2
Such compositions are economical, thixo- tronic and less subject to syneresis than the	tems by rebalancing the cosolven The gels of Examples 2 to 5 m	t fraction. Ly be mad	le
Such compositions are economical, thixo- tropic and less subject to syneresis than the brilliantine type compositions having a metal	The gels of Examples 2 to 5 mm by heating the oily solvent and the	t fraction. Ly be made e cosolver	le nt
Such compositions are economical, thixo- tropic and less subject to syneresis than the brilliantine type compositions having a metal soap, such as aluminium stearate, as the	The gels of Examples 2 to 5 m by heating the oily solvent and th to slightly above the melting p	t fraction.  Ly be made cosolver oint of the cosolver of the c	le nt ne
Such compositions are economical, thixo- tropic and less subject to syneresis than the brilliantine type compositions having a metal soap, such as aluminium stearate, as the 40 gelling agent, in mineral oil.	tems by rebalancing the cosolven. The gels of Examples 2 to 5 ms by heating the oily solvent and the to slightly above the melting p polyamide which is then introduce	t fraction.  Ly be made  e cosolver  oint of the  ed into the	le nt ne ne 105
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Such compositions are economical, thixotropic and less subject to syneresis than the brilliantine type compositions having a metal soap, such as aluminium stearate, as the 40 gelling agent, in mineral oil.  It is an important advantage of the invention that crystal clear gels can be made, although opaque or cloudy gels are also within the broader scope of the invention. Such 5 gels, whether crystal clear or non-clear, provide a thixotropic, non-greasy solid gel, which is liquefied quickly to a fluid by rubbing and is thus readily spread on and throughout the hair leaving the hair constitution of the spread on the state of the spread on the spread of the spread on th	tems by rebalancing the cosolven. The gels of Examples 2 to 5 ms by heating the oily solvent and the to slightly above the melting period to be solvents with a gitation until homogeneous. The temperatured to below 100°C and various are then introduced into the marring. It is then cured and filled it containers.  A further improvement has be by using appropriate mixtures of the resin and mineral oil. This crease of the mineral oil content sponding decrease in cost, while higher temperature stability and	t fraction.  y be made e cosolver oint of the ed into the ill the man ure is r us additive s with sti nto suitab en attained f cosolver s allows i with corr preserving preserving	de de la companya de
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	The mineral oil can be used in an amount from 1-80% by weight of the gel, and the polyamide in an amount from 1-40% by	curing our gelled compositions are protected against syneresis at considerably higher temperatures.	
5	weight, (advantageously 2-10% by weight) but in general the use of higher proportions of oil leads to some deterioration of the gel	The combination of the polyamide resin and oil with cosolvent can also be used for its extraordinary hair grooming and fixative effect in other than gel form. Examples of such are:	70
	bient temperatures, or higher, even the com-	Example 8—Spray Aerosol % By Weight Polyamide 8000 average MW 5.00	75
10	becomes evident, even though the product	Ethyl alcohol (specially denatured alcohol No. 40 anhydrous) 80.00 35%	
15	properties, the customer may assume that is is "spoiled." In such cases, special precau-	Perfume  Lanolin cil Propylene glycol mono laurata 13.40 0.60	80
	gelled thixotropic hair groom composition	Propellant 11 (Trichloro mono-fluoro methans) 50.00	85-
20.	after long periods of storage at amounts of peratures, can be provided. This is attained peratures, can be provided.	Propellant 12 (Dichloro di-fluoro methano) 50.00	•
25	by including in a stabilizer, an amide having a chain of at least 10 carbon atoms in its molecule. The following amides are advantageous as	The above is an example of the case men-	90
	stabilisers: — Lauryl diethanolamide Stearyl diethanolamide	mono-laurate acts as the liquid, oily non-	
30	Phenol Adduct	stead of lanolin oil, other lanolin derivatives which are compatible with the system may	73
	Linoleyi diethanolamide Coconut oil fatty acids diethanolamide Coconut oil diethanolamide	Example 9 — Clear Liquid Brilliantme Thixotropic Liquid type % By Weight Polyamide 8000 average MW 5.00	100
35	Tallow fatty acid diethanolamide Soy bean fatty acid diethanolamide Kritchevsky Condensates such as:  Coconut Oil-Diethanolamine	Propylene glycol mono laurate Light mineral oil Perfume 1.00	
40	Condensate and Lauric Acid-Diethanolamine	Free Flowing Liquid type Polyamide 8000 average MW Oleic Acid 10.00	105
	Condensate.  (Kritchevsky Condensates are products of the condensation of polyalkylol amines with fatty acids or glycerides thereof, said acids fatty acids or glycerides thereof, said acids	Perfume  Example No. 10 — Alcoholic Liquid Hair	r 110
45	cules).	Groom Parts by Weigh Polynomids 2000 average MW 2.00	1
•	such stabilisers the following example is given:	Light mineral oil Propylene glycol mono laurate  43.76 14.00	115
ж.	% By Weight Polyamide 8000 average MW 5.00	Ethyl alcohol (specially denatured alcohol No. 40.	.•
\$	Propylene giyool mono laurate 5.70 Lauryl diethanolamide 5.70 Oleic acid 10.90	Perfume 1.00  Example 11 — Emulsified Cream Hair	120 r
	Light mineral oil 64.00 Perfume 1.00	Propylene glycol mono laurate 15.00 Propylene glycol mono laurate 15.00 Propylene glycol mono laurate 10.00	125
61	Protection against syneresis can be gained	Triethanolamine 2300 Water 67.00 1.00	
6	or extended by curing the gel at a constant temperature between the melting point of the gel and ambient temperature. After such	In the above example, propylene given	n i, <b>i3</b> 0

oily, non-polar solvent as in Example 8.

The above emulsion is an example of an anionic type system. To those skilled in the art of emulsion making, it is readily under5 stood that similar systems can be produced using non-ionic and cationic emulsifiers or combinations of both.

The use of the above formulations, comprising the polyamide resin together with the 10 other ingredients, results in more enhanced hair grooming efficiency and produces a greater brilliance than that obtained with formulations of the conventional type.

Hair grooming properties of polyamides 15 in gel systems were substantiated by halfhead experiments, as described below:—

A gob of a clear gel about the size of a finger nail was weighed and liquefied by rubbing in the palm of the hand and applied to hair on half of one's head. An equivalent weight of the same formulation, but without the polyamide contained therein, was applied to the other half of the same head. Both halves were compared identically and the halves were compared for brilliance, grooming qualities, and hair fixative properties. The results indicated that the half-head containing the polyamide was superior in the aforementioned qualities. The preceding test was more demonstrative when hair switches of identical hair were used in place of the half-heads.

It should be recognised that in addition 35 to improving hair grooming properties, the use of the polyamide resins, as a gelling agent for solid brilliantines, is also new.

WHAT WE CLAIM IS:

 A hair grooming composition comprising a polyamide material at least partly dissolved in a liquid, oily, non-polar solvent consisting of or including a substance having a chain of at least 10 carbon atoms in its molecule, the polyamide material being a reaction product of an aliphatic polycarboxylic acid and an alkylene polyamine and having an average molecular weight between 2000 and 14000.

 A hair grooming composition as
 claimed in claim i in which the polyamide material is solid at ambient temperatures.

3. A hair grooming composition as claimed in claim 2 including a cosolvent which forms with the polyamide a solution 55 which is miscible with the liquid, oily solvent.

4. A hair grooming composition as claimed in claim 2 or 3 in which the liquid,

oily solvent has a viscosity between 50 and 70 Saybolt.

5. A hair grooming composition as claimed in claim 2, 3 or 4 which contains from 1% to 40% by weight of the polyamide material and from 1% to 80% by weight of the liquid oily solvent.

6. A hair grooming composition as claimed in claim 2, 3, 4 or 5 which contains from 2% to 10% by weight of the polyamide

7. A hair grooming composition as 70 claimed in any of the claims 2 to 6 in which there is incorporated a stabiliser which is an amide having a chain of at least 10 carbon atoms in its molecule.

8. A hair grooming composition as 75 claimed in claim 7 and which has been cured at a temperature between the melting point of the composition and ambient temperature.

9. A hair grooming composition as claimed in any one of claims 2 to 8 in which 80 the said polycarboxylic acid is polymerized linoleic acid.

10. A hair grooming composition as claimed in any one of claims 2 to 9 in which the average molecular weight of the polyamide material is from 5000 to 10000.

11. A hair grooming composition as claimed in claim 7 or 8 in which the said stabiliser is a diethanolamide having 12 to 18 carbon atoms in its molecule.

12. A hair grooming composition as claimed in any one of the preceding claims including a perfume.

13. A method of grooming hair comprising the step of applying a hair grooming 95 composition according to any one of the preceding claims to the hair.

14. A method of preparing a hair grooming composition according to claim 1 comprising the steps of forming a mixture of the polyamide material in the oily liquid non-polar solvent, and at least partially dissolving said polyamide in the solvent by applying heat to said solvent either before or after addition of said polyamide.

15. A hair grooming composition substantially as described in any one of the examples hereinbefore set forth.

16. A method of preparing a hair grooming composition substantially as herein-110 before described.

For the Applicants,
WILSON, GUNN & ELLIS,
Chartered Patent Agents,
57 Market Street,
Manchester, 1.

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